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### IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Application of:

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Nyce, J. W.

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P66 39255

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Examiner:

Hauda, K. M.

For:

AGENT, COMPOSITION, KIT & METHOD FOR TREATMENT OF DISORDERS ASSOCIATED WITH BRONCHOCONSTRICTION, INCLUDING ASTEMA

#### **DECLARATION UNDER 37 CFR 1.132**

Assistant Commissioner for Patents Washington, D C 20231

Sir/Madam:

Jonathan W. Nyce, Ph. D., hereby declares as follows.

- (1) I am the sole inventor in the shove-identified application.
- (2) I have read the above-identified patent application and the Office Action of October 1, 1997, and understand its contents. Claims 1-34 were rejected because, in the examiner's view, the specification is not broadly enabling of methods for reducing or treating a respiratory disease or condition.

#### (3) Oblective

The present work was conducted to demonstrate that the present invention is broadly applicable to anti-sense oligonucleotides ("oligos") specific to the adenosine A<sub>1</sub> receptor mRNAs.

### (4) Development

The following experimental study was conducted by me, or under my supervision, to show that the method of the invention is broadly suitable for use with anti-sense oligos designed as taught by this application and targeted to adenosine A<sub>i</sub> receptor mRNAs.

Anti-sense Oligo I was disclosed in the above-identified patent application. For the present work, an additional anti-sense phosphorothicate oligo targeted to the adenosine A, receptor (Oligo II) was designed and tested, as described in the above-identified patent application. This anti-sense oligo was designed for therapy on a selected species as described in the above patent application and is generally specific for that species, unless the segment of the adenosine A, receptor mRNA of another species selected for treatment happens to have a similar sequence. The anti-sense oligo was prepared as described below, and tested in vivo in a rabbit model for respiratory diseases, including bronchoconstriction, inflammation and allergy. This animal model is widely recognized by the scientific community as appropriate for testing therapies which will then be applied to humans who have breathing difficulties and impeded lung airways, as is the case in asthma and other conditions, as described in the above-identified application.

#### (5) Methods

#### (a) Anti-sense DNA

One oligo and its therapeutic effect were studied in a rabbit model and the results of these studies are reported and discussed below. This oligo was selected for this study to complement the data on SEQ ID NO: 1 (Oligo I), which is anti-sense to the adenosine A<sub>1</sub> receptor mRNA, provided in the above-identified patent application. The oligos, which are anti-sense to the adenosine A<sub>1</sub> receptor mRNA are identified as anti-sense Oligo I (SEQ ID NO: 1) and Oligo II (new oligo), which is a fragment targeted to a different region of the adenosine A<sub>1</sub> receptor mRNA. The design and synthesis of these anti-sense oligos was performed in accordance with the teachings of the above-identified patent application, particularly of Example 1.

(I) Anti-sense Oligo I The above-identified application disclosed anti-sense oligonucleotide I to the human A<sub>1</sub> adenosine receptor mRNA (EPI 2010, SEQ. ID NO: 1). Anti-sense oligo I is 21 nucleotide long, overlaps the initiation codon, and has the following sequence.

#### 5'- GAT GGA GGG CGG CAT GGC GGG -3'

The oligo I was previously shown to abrugate the adenosine-induced bronchoconstriction in allergic rabbits, and to reduce allergen-induced airway obstruction and bronchial hyperresponsiveness (BHR). See,

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Nyce, J. W. & Metzger, W. J., Nature, 385:721 (1977), a copy of which is enclosed.

(II) Anti-sense Oligo II: A phosphorothioate anti-sense oligo (EPI 2014) was designed in accordance with the invention to target the rabbit adenosine A<sub>1</sub> receptor mRNA region +936 to +956 relative to the initiation codon (start site). The anti-sense oligo II is 21 nucleotide long, and has the following sequence.

#### 5'-CTC GTC GCC GTC GCC GGC GGG-3'

(III) A<sub>1</sub> Mismatch Oligos: Two different mismatched oligonucleotides having the following sequences were used as controls for anti-sense oligo I (SEQ. ID NO: 1) described in (a) above.

A, MM 5'-GTA GGT GGC GGG CAA GGC GGG-3'
A, MM2 5'-GAT GGA GGC GGG CAT GGC GGG-3'

Anti-sense oligo I and the two mismatch anti-sense oligos had identical base content and general sequence structure. Homology searches in GENBANK (release 85.0) and EMBL (release 40.0) indicated that the anti-sense oligo I was specific, not only for the human, but also for the rabbit, adenosine A receptor genes, and that the mismatched controls were not candidates for hybridization with any known human or animal gene sequence.

(IV) Controls: Having established that mismatch-treated animals were equivalent to saline-treated animals, saline was used as the control while for Anti-sense Oligo II, control rabbits were administered 5.0 ml aerosolized sterile saline following the same schedule as for the anti-sense oligos in (II), (III), and (IV) above.

#### (b) Synthesis of Anti-sense Oligos

Phosphorothinate anti-sense oligos having the sequences described in (a) above, were synthesized on an Applied Biosystems Model 396 Oligonucleotide Synthesizer, and purified using NENSORB chromatography (DuPont, DE) TETD (tetraethylthiuram disulfide) was used as the sulfurizing agent during the synthesis. Anti-sense oligonucleotide II (EPI 2014) was synthesized and purified in this manner.

#### (c) Preparation of Allergic Rabbits

Neonatal New Zealand white Pasturella-free rabbits were immunized intraperitoneally within 24 hours of birth with 0.5 ml of 312 antigen units/ml house dust mite (D. farinze) extract (Berkeley Blologicals, Berkeley, CA) mixed with 10% keolin as

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previously described (Metzger, W. J. In: Late Phase Allergic Reactions, (Dorsch, W., Ed.), CRC Handbook, pp 347-362. CRC Press, Boca Raton, 1990; Ali, S., Metzger, W. J. and Mustafa, S. J., Am. J. Resp. Crit. Care Med. 149, 908, 1994). Immunizations were repeated weekly for the first month and then biweekly until the age of 4 months. These rabbits preferentially produce allergen specific IgE antibody, typically respond to aeroallergen challenge with both an early and late-phase asthmatic response, and show bronchial hyper responsiveness (BHR). Monthly intraperitoneal administration of allergen (312 units dust mite allergen, as above) continues to stimulate and maintain allergen specific IgE antibody and BHR. At 4 months of age, sensitized rabbits were prepared for aerosol administration as described by Ati et al. (199\_) (Ali, S., Metzger, W. J. and Mustafa, S. J., Am. J. Resp. Crit, Care Med. 149 (1994)).

#### (d) Dose-response Studies

#### (i) Experimental Setup

Aerosols of either adenosine (0-20 mg/ml), or anti sense or one of two mismatch oligonucleotides (5 mg/ml) were separately prepared with an ultrasonic nebulizer (Model 646, DeVilbiss, Somerset, PA), which produced acrosol droplets, 80% of which were smaller than \$\mu\$m in diameter. Equal volumes of the aerosols were administered directly to the lungs via an intratracheal tube.

The animals were randomized, and administered aerosolized adenosine. Day 1 pretreatment values for sensitivity to adenosine were calculated as the dose of adenosine causing a 50% loss of compliance ( $PC_{\infty}$  Adenosine). The animals were then administered either the serosolized anti-sense or one of the mismatch anti-sense oligos via an intratracheal tube (5 mg/1.0 ml), for 2 minutes, twice daily for 2 days (total dose, 20 mg). Post-treatment  $PC_{\infty}$ values were recorded (post-treatment challenge) on the morning of the third day. The results of these studies are provided in (6)(a)(iii) below.

#### (ii) Cross-over Experiments

For some experiments utilizing anti-sense Oligo I (SEQ ID NO: 1) and a corresponding mismatch oligonucleotide A<sub>1</sub>MM (Control), following a 2 week interval, the animals were crossed over, with those previously administered the mismatch control A<sub>1</sub>MM,

now receiving the anti-sense Oligo I, and those previously treated with the anti-sense oligo.

I, now receiving the mismatch control AMM oligo.

The number of animals per group was as follows. For mismatch AMM (Control 1), n=7, since one animal was lost in the second control arm of the experiment due to technical difficulties, for mismatch A<sub>1</sub>MM2 n=4 (Control 2) and for anti-sense Oliga I, n=8. The A<sub>2</sub>MM2 oligo treated animals were analyzed separately and were not part of the cross-over experiment. The treatment methods and measurements employed following the cross-over were identical to those employed in the first arm of the experiment.

In 6 of the 8 animals treated with the anti-sense Oligo I (SEQ. ID NO: 1), no PC<sub>30</sub> value could be obtained for adenosine doses of up to 20 mg/ml, which is the limit of solubility of adenosine. Accordingly, the PC<sub>30</sub> values for these animals were assumed to be 20 mg/ml for calculation purposes. The values given, therefore, represent a minimum figure for the effectiveness of the anti-sense oligonucleotides of the invention. Other groups of allergic rabbits (n=4 for each group) were administered 0.5 or 0.05 mg doses of the anti-sense oligo I (SEQ ID NO: 1), or the AMM Control Oligo in the manner and according to the schedule described above (the total doses being 2.0 or 0.2 mg). The results of these studies are provided in (6)(a)(iv) below.

#### (e) Anti-sense Oligo Formulation

Each one of the anti-sense oligos were separately solubilized in an aqueous solution and administered as described for anti-sense oligo I in (e) above, in four 5 mg aliquots (20 mg total dose) by means of a nebulizer via endotracheal tube, as described above.

The results obtained for ami-sense oligo I and its mismatch controls confirmed that the mismatch controls are equivalent to saline. See, Table 1 of Nyce & Metzger, Nature 385, 721-725, 1997. Because of this finding, saline was used as a control for pulmonary function studies employing anti-sense oligos II, III and IV.

# (f) Specificity of Otigo 1 for Adenosine A, Receptor (Receptor Binding Studies)

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Tissue from airway smooth muscle was dissected to primary, secondary and tertiary, bronchi from rabbits which had been administered 20 mg Oligo I (SEQ ID NO: 1; EPI 2010) in 4 divided doses over a period of 48 hours as described above. A membraue fraction was prepared according to the method of Ali et al. See, Ali, S., et al., Am. J. Resp. Crit. Care Med. 149,: 908 (1994).

The protein content was determined by the method of Bradford and plasma membranes were incubated with 0.2 U/ml adenosine deaminase for 30 minutes at 37°C to remove endogenous adenosine. See, Bradford, M. M. Anal, Biochem. 72, 240-254 (1976). The binding of [H]DPCPX, [H]NPC17731, or [H]CGS-21680 was measured as described by Jarvis et al. See, Jarvis, M.F., et al., Pharmacol. Exptl. Ther. 251, 888-893 (1989). The results of this study are shown in Table 1 and discussed in (6)(a)(ii) below.

#### Pulmonary Function Measurements **(g)** (Compliance CDYN and Resistance)

At 4 months of age, the immunized animals were anesthetized and relaxed with 1.5 ml of a mixture of ketamine HCl (35 mg/kg) and acepromazine maleute (1.5 mg/kg) administered intramuscularly. After induction of anesthesia, allergic rabbits were comfortably positioned supine on a soft molded animal board. Salve was applied to the eyes to prevent drying, and they were closed. The animals were then intubated with a 4.0 mm intermediate high-low cuffed Murphy 1 endotracheal tube (Mallinckrodt, Glen Falls, NY), as previously described by Zavala and Rhodes. See, Zavala and Rhodes, Proc. Soc. Exp. Biol. Med. 144; 509-512 (1973). A polyethylene catheter of OD 2.4 mm (Becton Dickinson, Clay Adams, Parsippany NJ) with an attached thin-walled latex balloon was passed into the esophagus and maintained at the same distance (approximately 16 cm) from the mouth throughout the experiment. The endotracheal tube was attached to a heated Fleisch pneumotach (size 00; DEM Medical, Richmond, VA), and the flow (v) measured using a Validyne differential pressure transducer (Model DP-45-16-1927, Validyne Engineering, Northridge, CA), driven by a Gould carrier amplifier (Model 11-4113, Gould Electronics, Cleveland, OH).

An exophageal halloon was attached to one side of the Validyne differential pressure transducer, and the other side was attached to the outflow of the endotracheal tube to obtain transpulmonary pressure (Pin). The flow was integrated to yield a continuous tidal volume, and the measurements of total lung resistance (R) and dynamic compliance (C<sub>600</sub>) were made

at isovolumetric and zero flow points. The flow, volume and pressure were recorded on an eight channel Gould 2000 W high frequency recorder and Cirn was calculated using the total volume and the difference in P at zero flow, and . R was calculated as the ratio of Ptp and V at midtidal lung volumes. These calculations were made automatically with the Buxco automated pulmonary mechanics respiratory analyzer (Model 6, Buxco Electronics, Sharon, CT), as previously described by Giles et al. See, Giles et al., Arch. Int. Pharmacodyn. Ther. 194: 213-232 (1971). The results obtained upon administration of oligo II on allergic rabbits are shown and discussed in (6)(b) below.

#### Measurement of Bronchial Hyperresponsiveness (BHR) **(b)**

Each allergic rabbit was administered histamine by serosol to determine their baseline hyperresponsiveness. Aerosols of either saline or histamine were generated using a DeVilbiss nebulizer (DeVilbiss, Somerset, PA) for 30 seconds and then for 2 minutes at each dose employed. The ultrasonic nebulizer produced aerosol droplets of which 80% were <5 micron in diameter. The histamine aerosol was administered in increasing concentrations (0.156 to 80 mg/ml) and measurements of pulmonary function were made after each dose. The BHR was then determined by calculating the concentration of histamine (mg/ml) required to reduce the C<sub>tro</sub> 50% from baseline (PC<sub>to Historius</sub>).

#### Cardiovascular Effect of Anti-sense Oilgo I (1)

The measurement of cardiac output and other cardiovascular parameters using Cardiomax utilizes the principal of thermal dilution in which the change in temperature of the block exiting the heart after a venous injection of a known volume of cool saline is monitured. A single rapid injection of cool saline was made into the right atrium via cannulation of the right jugular vein, and the corresponding changes in temperature of the mixed injectate and blood in the aprile arch were recorded via carmulation of the carutid artery by a temperature-sensing miniprobe.

Twelve hours after the allergic rabbits had been treated with acrosols of oligo I (EPI 2010; SEQ. ID NO: 1) as described in (d) above, the animals were anesthetized with 0.3 ml/kg of 80% Ketamine and 20% Xylazine. This time point coincides with previous data showing efficacy for SEQ. ID NO: 1. See, Nyce & Metzger, (1997). A thermocouple was then inserted into the left carotid artery of each rabbit, and was then advanced 6.5 cm and 1 15-09 ·10·193V

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secured with a silk ligature. The right jugular veln was then cannulated and a length of polyethylene tubing was inserted and secured.

A thermodilution curve was then established on a Cardiomax\*\* II (Columbus Instruments, Ohio) by injecting sterile saline at 20°C to determine the correctness of positioning of the thermocouple probe. After establishing the correctness of the position of the thermocouple, the femoral artery and vein were isolated. The femoral vein was used as a portal for drug injections, and the temoral artery for blood pressure and heart rate measurements. Once constant baseline cardiovascular parameters were established, Cardiomax\*\* measurements of blood pressure, heart rate, cardiac output, total peripheral resistance, and cardiac contractility were made.

## (J) Duration of Action of Oligo I (SEQ. ID NO: 1; EPI 2010)

Eight allergic rabbits received initially increasing log doses of adenosine by means of a nebulizer via an intra-tracheal tube as described in (f) above, beginning with 0.156 mg/ml until compliance was reduced by 50% (PC<sub>50 Advanter</sub>) to establish a baseline. Six of the rabbits then received four 5 mg aerosolized doses of (SEQ. ID NO: 1; EPI 2010) as described above. Two rabbits received equivalent amounts of saline vehicle as controls. Beginning 18 hours after the last treatment, the PC<sub>50 Advance</sub> values were tested again. After this point, the measurements were continued for all animals each day, for up to 10 days. The results of this study are shown in Figures 5 and 6 and discussed in (6)(a)(vii) below.

## (6) Results

#### (a) Anti-sense Oligo I

#### (i) Prior Work

The nucleotide sequence and other data for anti-sense Oligo I (SEQ. ID NO: 1), which is specific for the adenosine A<sub>1</sub> receptor, was provided in the original application. In addition, the application also contained experimental data showing the effectiveness of oligo I in down regulating the receptor number and activity.

Further information on anti-sense Oligo I was provided in a publication by my group. See, Nyce, J. W., and Metzger, W. J., Nature 385:721 (1997) (copy enclosed). The

Nyce & Metzger (1997) publication provided data showing that the anti-sense Oligo I (SEQ. ,-ID NO: 1):

- Reduces the number of adenosine A, receptors in the bronchial (1) smooth muscle of allergic rabbits in a dose dependent manner. See, Table 1 of Nyce & Metzger (1997).
- **(2)** Attenuates adenosine-induced bronchoconstriction and altergeninduced bronchoconstriction. See, Figure 4 of Nyce & Metzger (1997).
- Attenuates bronchial hyperresponsiveness as measured by PCso (3) histamine, a standard measurement to assess bronchial hyperresponsiveness. This result clearly demonstrates antiinflammatory activity of the anti-sense oligo I. See, Figure 4 of Nyce & Meizger (1997).
- (4) As expected, because it was designed to target it, is totally specific for the adenosine A, receptor, and has no effect at all at any dose on either the very closely related adenosine A2 receptor or the related bradykinin B2 receptor. See, Table 1 of Nyce & Metzger (1997), and Figure 2 accompanying this Declaration.
- (5)Mismatch control molecules (MMI and MM2; See, Figure 1 of Nyce & Metzger) had identical base composition and molecular weight but differed from the anti sense oligo I (SEQ ID NO: 1) by 6 and 2 mismatches, respectively. These mismatches, which are the minimum possible while still retaining identical base composition, produced absolutely no effect upon any of the targeted receptors (A., A<sub>2</sub> or B<sub>2</sub>). See, Figure 1 of Nyce & Metzger (1997).

These results, along with a complete lack of prior art on the use of anti-sense oligonucleotides, such as oligo I, targeted to the adenosine A, receptor, show the unexpected results obtain by mc. More generally, the anti-sense oligonucleotides of the invention which are directed to adenosine receptor burg targets, particularly targets associated with asthma, are not only unobvious over the art at large, but have been broadly enabled by the prior work reported in the above-identified application, the work reported in Nyce & Metzger (1997), and the further work reported here. These collective showings clearly enable and show the effectiveness, for their intended use, of the claimed agent and method for reducing or treating brunchoconstriction and lung inflammation.

#### (II) Oligo I Significantly Reduces Response to Adenosine Challenge

The receptor hinding experiment is described in (5)(f), and the results shown in Figures 1 and 2 accompanying this Declaration, and in Table 1 below which shows the

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binding characteristics of the adenosine A<sub>i</sub>-selective ligand [H]DPCPX and the bradykinine B, selective ligand [3HJNPC 17731 in membranes isolated from airway smooth muscle of A, adenosine receptor and B, brudykinin receptor anti-sense- and mismatch-treated allergic rabbits.

Table 1: Binding Characteristics of Three Anti-Sense Oligos

A <sub>1</sub> receptor		B <sub>2</sub> receptor		
Kd	B <sub>met</sub>	Kd	Bmax	
0.36±0.029 nM	19±1.52 Imoles•	0.39±0.031 nM	14.8±0.99 tmoles	
0.38±0.030 nM	32±2.56 fmoles*	0.41±0.028 nM	15.5±1.08 (moles	
0.37±0.030 nM	49±3.43 finoles	0.34±0.024 nM	15.0±1.06 fmoles	
0.34±0.027 nM	52.0±3.64	0.35±0.024 nM	14.0±1.0 fmoles	
0.37±0.033 nM	51 8=3.88	0.38±0.028 nM	14.6±1.02 fmoles	
	0.36±0.029 nM 0.38±0.030 nM 0.37±0.030 nM 0.34±0.027 nM	Kd     B <sub>max</sub> 0.36±0.029 nM     19±1.52 Imoles*       0.38±0.030 nM     32±2.56 fmoles*       0.37±0.030 nM     49±3.43 fmoles       0.34±0.027 nM     52.0±3.64	Kd     B <sub>max</sub> Kd       0.36±0.029 nM     19±1.52 Imoles*     0.39±0.031 nM       0.38±0.030 nM     32±2.56 fmoles*     0.41±0.028 nM       0.37±0.030 nM     49±3.43 fmoles     0.34±0.024 nM       0.34±0.027 nM     52.0±3.64     0.35±0.024 nM	

Refers to total oligo administered in four equivalently divided doses over a 48 hour period. Treatments and analyses were performed as described in methods. Significance was determined by repeated-measures analysis of variance (ANOVA), and Tukey's protected t test. n = 4-6 for all groups.

#### (ili) Dose-response Effect of Oligo I

Anti-sense Oligo I (SEQ ID NO: 1) was found to reduce the effect of adenosine administration to the animal in a dose-dependent manner over the dose range tested as shown in Table 2 below and in Figure 2.

<sup>\*</sup> Significantly different from mismatch control- and saline-treated groups, p<0.001;

<sup>\*\*</sup>Significantly different from mismatch control- and saline treated groups, p<0.05.

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Table 2: Dose-Response Effect to Anti-sense Oligo I

Total Dose	PC <sub>38 Addition</sub>			
(mg)	(mg Adenosine)			
Anti-sense Oligo I				
0.2	8.32 ±7.2			
2.0	14.0±7.2			
20	19.5±0.34			
A <sub>1</sub> MM oligo (control)				
0.2	2.51±0.46			
2.0	$3.13 \pm 0.71$			
20	$3.25 \pm 0.34$	$3.25 \pm 0.34$		

The above results were found to be gratismissly different under the Student's paired t test, p-0.05

Figure 2 shows that Ollgo I, an anti-administration A, receptor oligo, acts specifically on the adenosine A, receptor, but not on the adenosine A, receptor. These results stem from the treatment of rabbits with anti-sense oligo I or mismatch control oligo as described in (5) (d)(i) above and in Nyce & Metzger (1997) (four doses of 5 mg spaced 8 to 12 hours apart via nebulizer via endotracheal tube), bronchial smooth muscle tissue excised and the number of adenosine A<sub>1</sub> and adenosine A<sub>2</sub> receptors determined as reported in Nyce & Metzger (1997).

## (iv) Specificity of Oligo I for Target Gene Product

Oligo 1 is specific for the adenosine A<sub>i</sub> receptor whereas its mismatch controls had no activity. Figure 1 depicts the results obtained from the cross-over experiment described in (5)(d)(ii) above and in Nyce & Metrger (1997). As may be seen from the top and lower panels of Figure 1, the two mismatch controls evidence no effect on the PC<sub>N Adenosise</sub> value. On the contrary, the administration of anti-sense Oligo I (SEQ. ID NO: 1; EPI 2040) shows a seven-fold increase in the PCSI Administration value. The results shown in Figure 1 above clearly

indicate that anti sense Oligo I (SEQ. ID NO: 1; EPI 2010) reduces the response (attenuates the sensitivity) to exogenously administered adenosine when compared with a saline control. The results provided in Table 2 above clearly establish that the effect of the anti-sense oligo I is dose dependent (see, column 3 of Table 1).

Oligo I was also shown to be totally specific for the adenosine A, receptor, (see, top 3 rows), inducing no activity at either the closely related adenosine A, receptor (see, Figure 2, right hand panel), or at the bradykinin B, receptor (data not shown).

In addition, the results shown in Table 2 and Figure 2 establish that anti-sense oligo I decreases sensitivity to adenosine in a dose dependent manner, and that it does this in an anti-sense-dependent manner since neither of two mismatch control oligonucleotides show any effect on PC<sub>20 Adenosise</sub> values or on the number of adenosine A<sub>1</sub> receptors.

#### (v) Effect on Aeroallergen-induced Bronchoconstriction & Inflammation

Oligo I was shown to significantly reduce the histamine-induced effect in the rabbit model when compared to the mismatch oligos. Figure 3 shows the effect of anti-sense Oligo I and the mismatch oligos on allergen-induced airway obstruction and bronchial hypotresponsiveness in allergic rabbits. Pauels (a), (b), (c) and (d) represent the following.

Panel (a) shows the effect of anti-sense Oligo I (AAS; SEQ. ID NO:1) on allergen-induced airway obstruction. As calculated from the area under the curve, the anti-sense oligo I significantly inhibited allergen-induced airway obstruction (55%, p<0.05; repeated measures ANOVA, and Tukey's t test). Compare with panel (b) for mismatch AMM oligo (control).

Panel (b) shows the lack of effect of the mismatch Oligo A,MM (Control) on allergen induced airway obstruction.

Panel (c) shows the effect of the anti-sense Oligo I (A<sub>i</sub>AS; SEQ. ID NO:1) on allergen-induced BHR. As calculated from the  $PC_{50~Hirmone}$  value, (A<sub>i</sub>AS), the anti-sense oligo I significantly inhibited allergen-induced BHR in allergic rabbles (61%, p<0.05; repeated measures ANOVA, Tukey's t test). Compare with Panel (d) for mismatch A<sub>i</sub>MM oligo (Control)

Panel (d) shows a lack of effect of the AMM Mismatch Control on allergen-induced BHR.

The results shown in Figure 3, panel (a), indicate that ami-sense Oligo I (SEQ. ID of NO: 1; EPI 2010) is effective to protect against aeroallergen-induced bronchoconstriction (house dust mite). In addition, anti-sense oligo I was also found to be a potent inhibitor of dust mite-induced bronchial hyper responsiveness, as shown by its effects upon histamine sensitivity (panel(c)), indicating anti-inflammatory activity for anti-sense Oligo I.

## (vi) Anti-sense Oligo I is Free of Deleterious Side Effects

Oligo I was shown to be free of side effects that might be toxic to the recipient. No changes in arterial blood pressure, cardiac output, stroke volume, heart rate, total peripheral resistance or heart contractility (dPdT) were observed following administration of 2.0 or 20 mg oligo I. Figure 4 shows the results of the measurement of cardiac output (CO), stroke volume (SV), mean arterial pressure (MAP), heart rate (HR), total peripheral resistance (TPR), and contractility (dPdT) with a Cardiomax apparatus (Columbus Instruments, Ohio).

These results evidence that Oligo I has no detrimental effect upon critical cardiovascular parameters. More particularly, this oligo does not cause hypotension. This finding is of particular importance because other phosphorothioate anti-sense oligonucleotides have been shown in the past to induce hypotension in some model systems. Furthermore, the adenosine A<sub>1</sub> receptor plays an important role in sinoatrial conduction within the heart. Attenuation of the adenosine A<sub>1</sub> receptor by anti-sense oligo I might be expected to result, therefore, in deletenous extrapulmonary activity in response to the downregulation of the receptor. This is not the case. The anti-sense oligo I does not produce any deleterious extrapulmonary effects and renders the administration of the low doses of the present anti-sense oligo free of unexpected, undesirable side effects.

This demonstrates that when oligo I is administered directly to the lung, it does not reach the heart in significant quantities to cause deleterious effects. This is in contrast to traditional admostne receptor antagonists like the ophylline which do escape the lung and can cause deleterious, even life-threatening effects outside the lung.

#### (vii) Long Lasting Effect of Oligo I

Oligo I evidenced a long lasting effect as evidenced by the PC<sub>30</sub> adenosine and Resistance values obtained upon its administration prior to adenosine challenge. Figures 5 and 6 show the values obtained.

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Figure 5 shows the duration of the effect, with respect to the  $PC_{SI}$  adenosine of antisense Oligo I when administered in four equal doses of 5 mg each by means of a nebulizer via an endotracheal tube, as described above. The effect of the agent is significant over days 1 to 8 after administration. When the effect of the anti-sense oligo I had disappeared, the animals were administered saline aerosols (controls), and the  $PC_{SO}$  Administration values for all animals were measured again. Saline-treated animals showed base line  $PC_{SO}$  adenosine values (n=6).

Figure 6 shows the duration of the effect (with respect to Resistance) for six allergic rabbits which were administered 20 mg of anti-sense Oligo I (SEQ. ID NO: 1) as described above, upon airway resistance measured as also described above. The mean calculated duration of effect was 8.3 easys for both  $PC_{50}$  adenosine (p < 0.05) and resistance (p < 0.05). These results show that anti-sense oligo I has an extremely long duration of action (6.8 days), which is completely unexpected when compared to literature results of anti-sense oligonucleotides targeting other mRNAs.

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## (b) Anti-sense Oligo II

Anti-sense Oligo II, targeted to a different region of the adenosine  $A_i$  receptor mRNA, was found to be highly active against the adenosine  $A_i$ -mediated effects. The results of the experiment are shown in Figure 7, which evidences the effect of anti-sense Oligo II (EPI 2014) upon compliance (top figure) and resistance (lower figure) values when 20 mg anti-sense oligo II were administered to allergic rabbits as described above, and compliance and resistance values measured following an administration of adenosine as described above in (5)(g). Significant at p < 0.05 using paired t test, compliance; p < 0.01 for resistance.

The results of Figure 7 show that anti-sense Oligo II, which targets the adenosine A<sub>1</sub> receptor, effectively maintains compliance and reduces resistance upon adenosine challenge.

#### (c) Conclusions

The work described and results discussed above indicate that the two oligonucleotides which are anti-sense to the adenosine A<sub>i</sub> receptor mRNA, designed in accordance with the teachings of the above identified application, were found to be highly effective at countering or reducing effects mediated by the receptors they are targeted to. That is, the two anti-sense oligos targeting an adenosine A<sub>i</sub> receptor mRNA were shown capable of countering the

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effect of exogenously administered adenosine which is mediated by the specific receptor they are targeted to.

In addition, the results presented also show that the administration of the present agents results in extremely low or non-existent deleterious side effects or toxicity.

This represents 100% success in providing agents that are highly effective and specific in the treatment of bronchoconstriction and/or inflammation. This invention is applicable in the same manner to all fragments which are anti-sense to adenosine A<sub>1</sub> receptor mRNAs.

These are clearly superior results which could not have been expected based on the knowledge of the art at the time of this invention. The experimental data and results provided are clearly enabling of the fragment oligonucleotides targeted to lung adenosine  $A_1$  receptors described and claimed in the above-identified application.

- (8) I hereby declare that all statements made herein of my own knowledge are true, and that all statements made on information and belief are believed to be true; and that these statements are made with the knowledge that willful false statements and the like are punishable by fine or imprisonment, or both, under § 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application of any patent issuing thereon.
  - (9) Declarant further sayth not.

Date

Yonathan W. Nyce, Ph. D

A: Ndec255.jwo

3/06/98

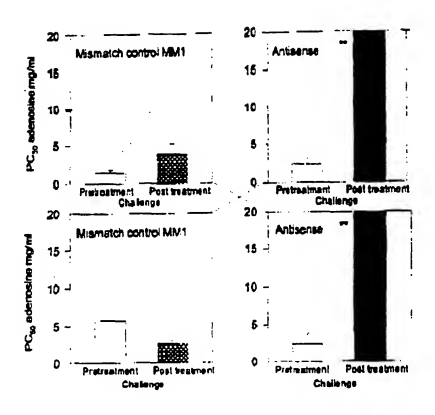


Figure 1: EPI 2010 Data Summary from both arms of crossover experiment PCs Adenosine A.AS

A. MM Control		At NEW 2 CONTUCT		Airo		
					Pre ODN	
	3 56 ± 1.02	$3.25 \pm 0.34$	$2.46 \pm 0.50$	2.81 ± 0.70	$2.36 \pm 0.68$	>19.5 ±0.34**

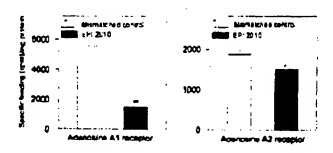


Figure 2.

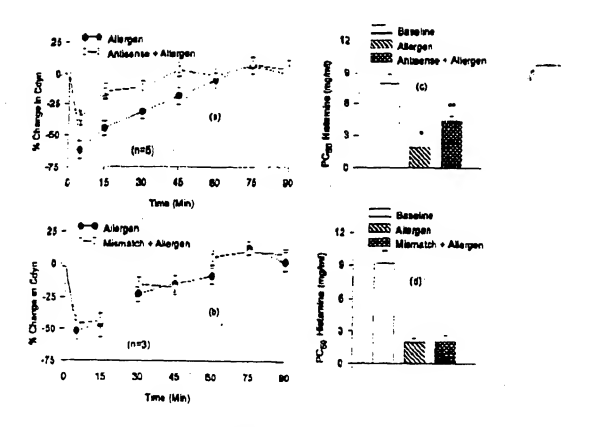


Figure 3.

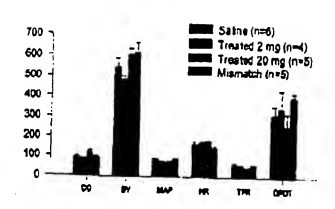


Figure 4.

# EPI 2010 Duration of Effect (n=6)

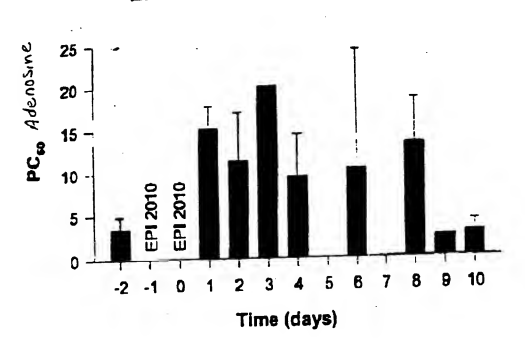
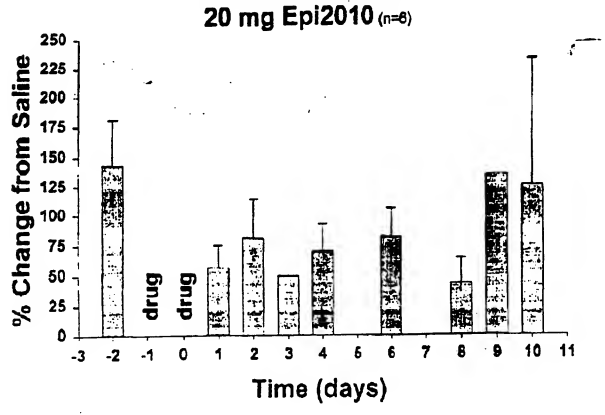
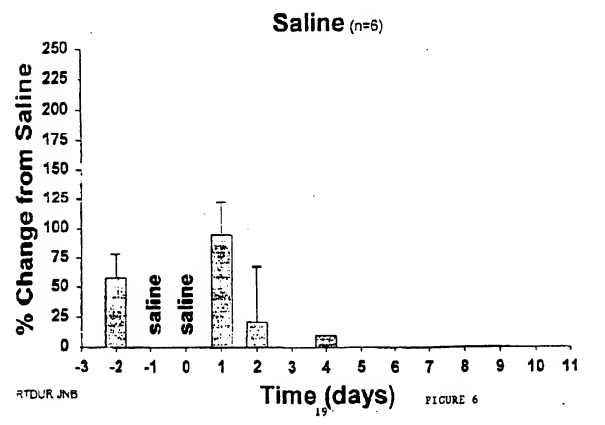


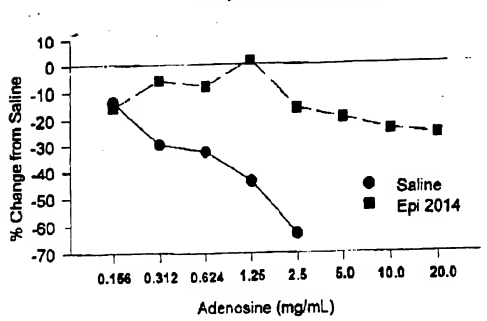
Figure 5.

# EpiGen sis Duration Study Resistance Changes





# Compliance, Epi 2014



# Resistance, Epi 2014

